		IS, MEDLINE,				ΑT	15:54:28	ON	28	AUG	2002
L13	3892	PHOSPHATIDY:	LINOSITO	L-4-PHOS	SPHATE						
L14	1001	5-KINASE									
Lİ5	652	L13 AND L14									
L16	268	EC 2.7.1.68									
L17	58	PIP 5-KINAS	Ε								
L18	74061	ANTISENSE									
L19	14625	RIBOZYME									
L20	4	L15 AND L18									
L21	4	DUP REM L20	(O DUPL	ICATES F	REMOVED)						
L22	2	L15 AND L19									
L23	0	L16 AND L18									
L24	0	L16 AND L19									
L25	0	L17 AND L18									
L26	0	L17 AND L19									

L12 ANSWER 1 OF 3 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR(S):

L12 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2002:369320 CAPLUS

136:381056

The contribution of 700,000 ORF sequence tags to the definition of the human transcriptome Camargo, Anamaria A.; Samaia, Helena P. B.; Dias-Neto, Emmanuel; Simao, Daniel F.; Migotto, Italo A.; Briones, Marcelo R. S.; Costa, Fernando F.; Nagai, Maria Aparecida; Verjovski-Almeida, Sergio; Zago, Marco A.; Andrade, Luis Eduardo C.; Carrer, Helaine; El-Dorry, Hamza F. A.; Espreafico, Enilza M.; Habr-Gama, Angelita; Giannella-Neto, Daniel; Goldman, Gustavo H.; Gruber, Arthur; Hackel, Christine; Kimura, Edna T.; Maciel, Rui M. B.; Marie, Suely K. N.; Martins, Elizabeth A. L.; Nobrega, Marina P.; Paco-Larson, Maria Luisa; Pardini, Maria Ines M. C.; Pereira, Goncalo G.; Pesquero, Joao Bosco; Rodrigues, Vanderlei; Rogatto, Silvia R.; Da Silva, Ismael D. C. G.; Sogayar, Mari C.; Sonati, Maria De Fatima; Tajara, Eloiza H.; Valentini, Sandro R.; Alberto, Fernando L.; Amaral, Maria Elisabete J.; Aneas, Ivy; Arnaldi, Liliane A. T.; De Assis, Angela M.; Bengtson, Mario Henrique; Bergamo, Nadia Aparecida; Bombonato, Vanessa; De Camargo, Maria E. R.; Canevari, Renata A.; Carraro, Dirce M.; Cerutti, Janete M.; Correa, Maria Lucia C.; Correa, Rosana F. R.; Costa, Maria Cristina R.; Curcio, Cyntia; Hokama, Paula O. M.; Ferreira, Ari J. S.; Furuzawa, Gilberto K.; Gushiken, Tsieko; Ho, Paulo L.; Kimura, Elza; Krieger, Jose E.; Leite, Luciana C. C.; Majumder, Paromita; Marins, Mozart; Marques, Everaldo R.; Melo, Analy S. A.; Melo, Monica; Mestriner, Carlos Alberto; Miracca, Elisabete C.; Miranda, Daniela C.; Nascimento, Ana Lucia T. O.; Nobrega, Francisco G.; Ojopi, Elida P. B.; Pandolfi, Jose Rodrigo C.; Pessoa, Luciana G.; Prevedel, Aline C.; Rahal, Paula; Rainho, Claudia A.; Reis, Eduardo M. R.; Ribeiro, Marcelo L.; Da Ros, Nancy; De Sa, Renata G.; Sales, Magaly M.; Sant'anna, Simone Cristina; Dos Santos, Mariana L.; Da Silva, Aline M.; Da Silva, Neusa P.; Silva, Wilson A., Jr.; Da Silveira, Rosana A.; Sousa, Josane F.; Stecconi, Daniella; Tsukumo, Fernando; Valente, Valeria; Soares, Fernando; Moreira, Eloisa S.; Nunes, Diana N.; Correa, Ricardo G.; Zalcberg, Heloisa; Carvalho, Alex F.; Reis, Luis F. L.; Brentani, Ricardo R.; Simpson, Andrew J. G.; De Souza, Sandro J. Ludwig Institute for Cancer Research, Sao Paulo, 01509-010, Brazil

CORPORATE SOURCE:

SOURCE:

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:

Proceedings of the National Academy of Sciences of the United States of America (2001), 98(21), 12103-12108

CODEN: PNASA6; ISSN: 0027-8424 National Academy of Sciences

Journal English

Open reading frame expressed sequences tags (ORESTES) differ from conventional ESTs by providing sequence data from the central protein coding portion of transcripts. A total of 696,745 ORESTES sequences were generated from 24 human tissues and a subset of the data that correspond to a set of 15,095 full-length mRNAs used as a means of assessing the efficiency of the strategy and its potential contribution to the definition of the human transcriptome. It was estd. that ORESTES sampled over 80% of all highly and moderately expressed, and between 40% and 50% of rarely expressed, human genes. In the most thoroughly sequenced tissue, the breast, the 130,000 ORESTES generated are derived from

transcripts from an estd. 70% of all genes expressed in that tissue, with an equally efficient representation of both highly and poorly expressed genes. In this respect, the capacity of the ORESTES strategy both for gene discovery and shotgun transcript sequence generation significantly exceeds that of conventional ESTs. The distribution of ORESTES is such that many human transcripts are now represented by a scaffold of partial sequences distributed along the length of each gene product. The exptl. joining of the scaffold components, by reverse transcription-PCR, represents a direct route to transcript finishing that may represent a useful alternative to full-length cDNA cloning. [This abstr. record is one of many records for this document necessitated by the large no. of index entries required to fully index the document and publication system constraints.].

L12 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:321987 CAPLUS

DOCUMENT NUMBER:

135:148100

TITLE:

Verification and initial annotation of the NIA mouse

15K cDNA clone set

AUTHOR(S):

Kargul, George J.; Dudekula, Dawood B.; Qian, Yong; Lim, Meng K.; Jaradat, Saied A.; Tanaka, Tetsuya S.;

Carter, Mark G.; Ko, Minoru S. H.

CORPORATE SOURCE:

Developmental Genomics and Aging Section, Laboratory

of Genetics, National Institute on Aging, National

Institutes of Health, Baltimore, MD, USA

SOURCE:

Nature Genetics (2001), 28(1), 17-18 CODEN: NGENEC; ISSN: 1061-4036

Nature America Inc.

PUBLISHER:
DOCUMENT TYPE:

Journal

LANGUAGE: English

A set of 15,247 unique oligo(dT)-primed cDNA clones (NIA Mouse 15K) based on 52,374 3'-expressed sequence tags (ESTs) has been made available for expression profiling in mouse models. To verify clone identity and obtain protein-coding information, the cDNA clones were resequenced, with an av. insert size of 1.5 kb, from both 3' and 5' ends (GenBank Accession Nos. BG062929-BG088954). Of 13,968 clones that were verifiable, 4.1% had correctable addressing errors or ambiguous addresses. Using all available sequence information to characterize the cDNA clone set indicated that up to 75% of mammalian genes in the public database are included in this set, and the remaining 3653 ESTs very likely represent novel genes that can be studied by expression profiling with the 15K microarray. All available information about individual cDNAs in this cDNA microarray are available through NCBI and MGD and in an ORACLE relational database accessible through the web site http://lgsun.grc.nia.nih.gov/cgi-bin/prol. abstr. record is one of 6 records for this document necessitated by the large no. of index entries required to fully index the document and publication system constraints.].

REFERENCE COUNT:

16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

2000:576531 CAPLUS

DOCUMENT NUMBER:

133:160369

TITLE:

Genome-wide expression profiling of mid-gestation placenta and embryo using a 15,000 mouse developmental

cDNA microarray

AUTHOR(S):

Tanaka, Tetsuya S.; Jaradat, Saied A.; Lim, Meng K.;
Kargul, George J.; Wang, Xiaohong; Grahovac, Marija
J.; Pantano, Serafino; Sano, Yuri; Piao, Yulan;
Nagaraja, Ramaiah; Doi, Hirofumi; Wood, William H.,

III; Becker, Kevin G.; Ko, Minoru S. H.

CORPORATE SOURCE:

Laboratory of Genetics, National Institutes of Health,

Baltimore, MD, 21224-5820, USA

SOURCE:

Proceedings of the National Academy of Sciences of the

United States of America (2000), 97(16), 9127-9132

CODEN: PNASA6; ISSN: 0027-8424 National Academy of Sciences

PUBLISHER: DOCUMENT TYPE:

Journal

English LANGUAGE:

Complementary DNA microarray technol. has been increasingly used to monitor global gene expression patterns in various tissues and cell types. However, applications to mammalian development have been hampered by the lack of appropriate cDNA collections, particularly for early developmental stages. To overcome this problem, a PCR-based cDNA library construction method was used to derive 52,374 expressed sequence tags from pre- and peri-implantation embryos, embryonic day (E) 12.5 female gonad/mesonephros, and newborn ovary. From these cDNA collections, a microarray representing 15,264 unique genes (78% novel and 22% known) was assembled. In initial applications, the divergence of placental and embryonic gene expression profiles was assessed. At stage E12.5 of development, based on triplicate expts., 720 genes (6.5%) displayed statistically significant differences in expression between placenta and embryo. Among 289 more highly expressed in placenta, 61 placenta-specific genes encoded, for example, a novel prolactin-like protein. The no. of genes highly expressed (and frequently specific) for placenta has thereby been increased 5-fold over the total previously reported, illustrating the potential of the microarrays for tissue-specific gene discovery and anal. of mammalian developmental programs. The sequences of the expressed sequence tags are available in the GenBank database at Accession Nos. AW537829-AW545916, AW535144-AW537732, AW545922-AW559162, and AF272368. [This abstr. record is one of 6 records for this document necessitated by the large no. of index entries required to fully index the document and publication system constraints.].

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L20 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2002:539817 CAPLUS 137:90191 DOCUMENT NUMBER: TITLE:

Identification, cloning, characterization and

therapeutic use of a human phosphatidylinositol 4phosphate 5-kinase family

member 56634

INVENTOR(S):

Meyers, Rachel A.; Rudolph-Owen, Laura A. Millennium Pharmaceuticals, Inc., USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 122 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE KIND DATE PATENT NO. \_\_\_\_\_ WO 2002055682 A2 20020718 WO 2001-US47782 20011113 2002055682 A2 20020718 WO 2001-US47782 20011113

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG APPLN. INFO:: US 2000-248325P P 20001114

PRIORITY APPLN. INFO.:

The invention provides isolated nucleic acids mols., designated 56634 nucleic acid mols., which encode novel phosphatidylinositol

4-phosphate 5-kinase members. The

cDNA sequence and the encoded amino acid sequence of a human

phosphatidylinositol 4-phosphate 5-

kinase homolog 56634 (clone Fbh56634FL) are disclosed. Tissue-specific expression profiles, and structural motifs of the polypeptide are provided. The invention also provides antisense nucleic acid mols., recombinant expression vectors contg. 56634 nucleic acid mols., host cells into which the expression vectors have been introduced, and nonhuman transgenic animals in which a 56634 gene has been introduced or disrupted. The invention still further provides isolated 56634 proteins, fusion proteins, antigenic peptides and anti-56634 antibodies. Diagnostic methods utilizing compns. of the invention are also provided.

L20 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2002:521952 CAPLUS

DOCUMENT NUMBER:

137:74468

TITLE:

Human phosphatidylinositol-4-

phosphate 5-kinase and

cDNA and drug screening targeted to regulation and other therapeutic application for related diseases

INVENTOR(S):

Zhu, Zhimin

PATENT ASSIGNEE(S):

Bayer Aktiengesellschaft, Germany

SOURCE:

PCT Int. Appl., 127 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. \_\_\_\_\_\_

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WO 2001-EP15321 20011227
    WO 2002053714
                     A2
                          20020711
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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                                      US 2001-259217P P 20010103
PRIORITY APPLN. INFO.:
                                      US 2001-331474P P 20011116
    A human phosphatidylinositol-4-phosphate
    5-kinase and cDNA and sequence homologs thereof are
    disclosed. The mRNA expression profile in various human tissues is
    provided. Methods for expressing and prepg. related products using
    recombinant cells are described. These recombinant cells, the enzyme, or
    nucleic acids encoding the enzyme are useful in screening for modulators
    of the enzymic activity or gene expression. Methods of screening for its
    modulators and using them for the treatment of various disease and their
    effectiveness (in vivo testing of compds./target validation) are
    described. Reagents that regulate human phosphatidylinositol-
    4-phosphate 5-kinase and reagents
    which bind to human phosphatidylinositol-4-
    phosphate 5-kinase gene products can play a
    role in preventing, ameliorating, or correcting dysfunctions or diseases
    including, but not limited to, cancer, asthma, and COPD.
L20 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
                        2001:816855 CAPLUS
                        135:353850
DOCUMENT NUMBER:
                        Protein and cDNA sequences of 13 kDa human
TITLE:
                        phosphatidylinositol-4-
                        phosphate-5-kinase
                        isoenzyme II .beta. subunit (PIPKII.beta.)-like
                        protein and therapeutic use thereof
                        Mao, Yumin; Xie, Yi
INVENTOR(S):
                        Shanghai Biowindow Gene Development Inc., Peop. Rep.
PATENT ASSIGNEE(S):
                        China
                        PCT Int. Appl., 35 pp.
SOURCE:
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
                        Chinese
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                 KIND DATE
                                         APPLICATION NO.
     PATENT NO.
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     WO 2001083688 A2
WO 2001083688 A3
                                         WO 2001-CN648
                                                          20010428
                           20011108
                           20020103
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         DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                       CN 2000-115550 20000429
                      A 20011114
     CN 1321756
                                       CN 2000-115550 A 20000429
PRIORITY APPLN. INFO.:
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The invention provides protein and cDNA sequences for 13 kDa novel human protein cloned from fetal brain, and which have similar expression pattern

with human PIPKII.beta.-like proteins (PIPKII.beta.). The invention also relates to constructing PIPKII.beta.-like protein gene expression vectors to prep. recombinant PIPKII.beta.-like protein using prokaryote or eukaryote cells. Methods of expressing and prepg. recombinant PIPKII.beta.-like protein and its antibody are described. Methods of using PIPKII.beta.-like protein or genes for the treatment of various kinds of diseases, such as cancer, blood diseases, HIV infection, immune diseases and inflammation are also disclosed.

L20 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1999:672845 CAPLUS

DOCUMENT NUMBER: 131:309275

TITLE: Nucleic acid sequences and proteins associated with

aging in human cells

INVENTOR(S): Burmer, Glenna C.; Brown, Joseph P. PATENT ASSIGNEE(S): Lifespan Biosciences, Inc., USA

SOURCE: PCT Int. Appl., 87 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO. DATE
      PATENT NO.
                          KIND DATE
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      WO 9952929 A1 19991021 WO 1999-US8314 19990415
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            MD, RU, TJ, TM

RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
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      AU 9935639
                                 A1
                                        19991101
                                                        EP 1999-917547
                                                                                     19990415
      EP 1071698
                                A1
                                        20010131
                 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                  IE, FI
                                                               JP 2000-543485
                                                                                        19990415
                                         20020416
       JP 2002511240
                                 T2
                                                          US 1998-81887P P 19980415
PRIORITY APPLN. INFO.:
                                                                                   A 19990414
                                                          US 1999-292758
                                                                                 W 19990415
                                                          WO 1999-US8314
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This invention relates to the discovery of nucleic acids assocd. with cell AB proliferation, cell cycle arrest, cell death, and aging-related diseases such as progeria and Werner syndrome. Such sequences can be used to det. the aging status of a cell population, e.g., whether a cell is aging or is undergoing senescence. Moreover, the present invention provides sequences indicative of the proliferation state or youth of a cell. In addn. the present invention provides sequences assocd. with the aging of skin cells and, in particular, fibroblast cells. The isolated nucleic acids can be used to det. the aging status of a cell population. In addnl., they can also be targeted and their level of expression altered by, for example, gene therapy methods. Such methods can be used to slow or stop the aging process of the cell population, to arrest the growth of a proliferating cell population such as a tumor cell population, to promote division in cells which are prematurely arrested, to det. that a cell population is healthy and rapidly dividing, and to det. that a cell population is not dividing and proliferating. Further, the present invention provides isolated nucleic acids assocd. with cyclin A.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L22 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2002 ACS 2002:521952 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER:

137:74468

TITLE:

Human phosphatidylinositol-4-

phosphate 5-kinase and

cDNA and drug screening targeted to regulation and other therapeutic application for related diseases

Zhu, Zhimin INVENTOR(S):

PATENT ASSIGNEE(S):

Bayer Aktiengesellschaft, Germany

SOURCE:

PCT Int. Appl., 127 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	ENT	NO.		KI	ND	DATE			A)	PPLI	CATI	ои ис	ο.	DATE			
WO	WO 2002053714					WO 2001-EP15321 20011227											
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		LS.	LT.	LU.	LV.	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	ΝZ,	OM,	PH,
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		BF.	ВJ.	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG
BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG PRIORITY APPLN. INFO.: US 2001-259217P P 20010103																	
21.201.22								1	US 2	001-	3314	74P	P	2001	1116		

## A human phosphatidylinositol-4-phosphate AB

5-kinase and cDNA and sequence homologs thereof are The mRNA expression profile in various human tissues is disclosed. provided. Methods for expressing and prepg. related products using recombinant cells are described. These recombinant cells, the enzyme, or nucleic acids encoding the enzyme are useful in screening for modulators of the enzymic activity or gene expression. Methods of screening for its modulators and using them for the treatment of various disease and their effectiveness (in vivo testing of compds./target validation) are described. Reagents that regulate human phosphatidylinositol-

4-phosphate 5-kinase and reagents

which bind to human phosphatidylinositol-4phosphate 5-kinase gene products can play a

role in preventing, ameliorating, or correcting dysfunctions or diseases including, but not limited to, cancer, asthma, and COPD.

L22 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1999:672845 CAPLUS

DOCUMENT NUMBER:

131:309275

TITLE:

SOURCE:

Nucleic acid sequences and proteins associated with

aging in human cells

INVENTOR(S): PATENT ASSIGNEE(S):

Burmer, Glenna C.; Brown, Joseph P. Lifespan Biosciences, Inc., USA

PCT Int. Appl., 87 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9952929	A1	19991021	WO 1999-US8314	19990415

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AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
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            CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
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This invention relates to the discovery of nucleic acids assocd. with cell AB proliferation, cell cycle arrest, cell death, and aging-related diseases such as progeria and Werner syndrome. Such sequences can be used to det. the aging status of a cell population, e.g., whether a cell is aging or is undergoing senescence. Moreover, the present invention provides sequences indicative of the proliferation state or youth of a cell. In addn. the present invention provides sequences assocd. with the aging of skin cells and, in particular, fibroblast cells. The isolated nucleic acids can be used to det. the aging status of a cell population. In addnl., they can also be targeted and their level of expression altered by, for example, gene therapy methods. Such methods can be used to slow or stop the aging process of the cell population, to arrest the growth of a proliferating cell population such as a tumor cell population, to promote division in cells which are prematurely arrested, to det. that a cell population is healthy and rapidly dividing, and to det. that a cell population is not dividing and proliferating. Further, the present invention provides isolated nucleic acids assocd. with cyclin A.

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The contribution of 700,000 ORF sequence tags to the definition of the human transcriptome Camargo, Anamaria A.; Samaia, Helena P. B.; Dias-Neto, Emmanuel; Simao, Daniel F.; Migotto, Italo A.; Briones, Marcelo R. S.; Costa, Fernando F.; Nagai, Maria Aparecida; Verjovski-Almeida, Sergio; Zago, Marco A.; Andrade, Luis Eduardo C.; Carrer, Helaine; El-Dorry, Hamza F. A.; Espreafico, Enilza M.; Habr-Gama, Angelita; Giannella-Neto, Daniel; Goldman, Gustavo H.; Gruber, Arthur; Hackel, Christine; Kimura, Edna T.; Maciel, Rui M. B.; Marie, Suely K. N.; Martins, Elizabeth A. L.; Nobrega, Marina P.; Paco-Larson, Maria Luisa; Pardini, Maria Ines M. C.; Pereira, Goncalo G.; Pesquero, Joao Bosco; Rodrigues, Vanderlei; Rogatto, Silvia R.; Da Silva, Ismael D. C. G.; Sogayar, Mari C.; Sonati, Maria De Fatima; Tajara, Eloiza H.; Valentini, Sandro R.; Alberto, Fernando L.; Amaral, Maria Elisabete J.; Aneas, Ivy; Arnaldi, Liliane A. T.; De Assis, Angela M.; Bengtson, Mario Henrique; Bergamo, Nadia Aparecida; Bombonato, Vanessa; De Camargo, Maria E. R.; Canevari, Renata A.; Carraro, Dirce M.; Cerutti, Janete M.; Correa, Maria Lucia C.; Correa, Rosana F. R.; Costa, Maria Cristina R.; Curcio, Cyntia; Hokama, Paula O. M.; Ferreira, Ari J. S.; Furuzawa, Gilberto K.; Gushiken, Tsieko; Ho, Paulo L.; Kimura, Elza; Krieger, Jose E.; Leite, Luciana C. C.; Majumder, Paromita; Marins, Mozart; Marques, Everaldo R.; Melo, Analy S. A.; Melo, Monica; Mestriner, Carlos Alberto; Miracca, Elisabete C.; Miranda, Daniela C.; Nascimento, Ana Lucia T. O.; Nobrega, Francisco G.; Ojopi, Elida P. B.; Pandolfi, Jose Rodrigo C.; Pessoa, Luciana G.; Prevedel, Aline C.; Rahal, Paula; Rainho, Claudia A.; Reis, Eduardo M. R.; Ribeiro, Marcelo L.; Da Ros, Nancy; De Sa, Renata G.; Sales, Magaly M.; Sant'anna, Simone Cristina; Dos Santos, Mariana L.; Da Silva, Aline M.; Da Silva, Neusa P.; Silva, Wilson A., Jr.; Da Silveira, Rosana A.; Sousa, Josane F.; Stecconi, Daniella; Tsukumo, Fernando; Valente, Valeria; Soares, Fernando; Moreira, Eloisa S.; Nunes, Diana N.; Correa, Ricardo G.; Zalcberg, Heloisa; Carvalho, Alex F.; Reis, Luis F. L.; Brentani, Ricardo R.; Simpson, Andrew J. G.; De Souza, Sandro J.

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Open reading frame expressed sequences tags (ORESTES) differ from conventional ESTs by providing sequence data from the central protein coding portion of transcripts. A total of 696,745 ORESTES sequences were generated from 24 human tissues and a subset of the data that correspond to a set of 15,095 full-length mRNAs used as a means of assessing the efficiency of the strategy and its potential contribution to the definition of the human transcriptome. It was estd. that ORESTES sampled over 80% of all highly and moderately expressed, and between 40% and 50% of rarely expressed, human genes. In the most thoroughly sequenced tissue, the breast, the 130,000 ORESTES generated are derived from

transcripts from an estd. 70% of all genes expressed in that tissue, with an equally efficient representation of both highly and poorly expressed genes. In this respect, the capacity of the ORESTES strategy both for gene discovery and shotgun transcript sequence generation significantly exceeds that of conventional ESTs. The distribution of ORESTES is such that many human transcripts are now represented by a scaffold of partial sequences distributed along the length of each gene product. The exptl. joining of the scaffold components, by reverse transcription-PCR, represents a direct route to transcript finishing that may represent a useful alternative to full-length cDNA cloning. [This abstr. record is one of many records for this document necessitated by the large no. of index entries required to fully index the document and publication system constraints.].